Amendments to the Claims:

This listing of claims will replace all prior versions, and listings, of claims in the application:

Listing of Claims:

- 1. (Currently amended) A method for detecting a target biopolymer in a sample, comprising:
 - (a) preparing a microarray of said sample by dispensing aliquots of said sample at discrete sites onto a substrate and immobilizing said target biopolymer on said substrate, wherein the microarray is an array of dots, each dot having a diameter from about 1 to 500 microns, wherein each of said aliquots contains the same amount of said target biopolymer;
 - (b) contacting said microarray with [a probe biopolymer] one or a plurality of probe biopolymers under conditions that allow the formation of [a complex] one or a plurality of complexes, each complex comprising said target biopolymer and [said probe biopolymer] one of said probe biopolymers, wherein said probe [biopolymer is] biopolymers are [applied to dots individually] deposited on said dots in said microarray; and
 - (c) detecting the presence of <u>and quantifying</u> said [complex] <u>complexes</u> as a measurement for the presence or the amount of the target biopolymer in said sample.

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2. (Original) The method of claim 1, wherein the preparation of said

microarray further comprises dispensing said sample aliquots on said substrate by

a method selected from the group consisting of jet printing, piezoelectric dispensing

methods, solenoid dispensing methods, thermal dispensing methods, solid pin

contact printing methods, capillary quill contact printing methods, microfluidic-

based printing, and silk screening.

3. (Original) The method of claim 1, wherein said aliquots comprise picomole

amounts of said target biopolymer.

4. (Original) The method of claim 1, wherein said aliquots comprise

femtomole amounts of said target biopolymer.

5. (Original) The method of claim 1, wherein said aliquots comprise attomole

amounts of said target biopolymer.

6. (Original) The method of claim 1, wherein said aliquots comprise

zeptomole amounts of said target biopolymer.

7. (Original) The method of claim 1, wherein said target biopolymer or said

probe biopolymer is selected from the group consisting of polynucleotides,

polypeptides, carbohydrates, and analogs thereof.

8. (Currently amended) The method of claim 7, wherein said polynucleotide

is selected from the group consisting of amplified DNA, cDNA, single-stranded

DNA, double-stranded DNA, peptide nucleic acids (PNA), RNA, and mRNA.

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9. (Original) The method of claim 7, wherein said polypeptide is selected

from the group consisting of antibodies, antibody fragments, antigens, ligands, and

receptors.

10. (Currently amended) The method of claim 1, wherein said target

biopolymer is a first polynucleotide and said probe biopolymer is a second

polynucleotide that is complementary to said [target] first polynucleotide.

11. (Original) The method of claim 1, wherein said target biopolymer is a

receptor and said probe biopolymer is a ligand for said receptor.

12. (Original) The method of claim 1, wherein said target biopolymer is an

antigen and said probe biopolymer is an antibody specific for said antigen.

13. (Original) The method of claim 1, wherein said probe is labeled with a

reporter selected from the group consisting of dyes, chemiluminescent compounds.

enzymes, fluorescent compounds, metal complexes, magnetic particles, biotin,

haptens, radio frequency transmitters, radioluminescent compounds, radioactive-

labeled biomolecules, dye-labeled beads, quantum dots, and bar coded particles.

14. (Original) The method of claim 1, wherein said substrate is made of

crosslinked polymers, porous foam, nitrocellulose, nylon, glass, silica, ceramic, gold,

porous metallic materials, non-porous metallic materials, and surface modified

materials.

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15. (Original) The method of claim 14, wherein said crosslinked polymers are

selected from the group consisting of polypropylene, polyethylene, polystyrene, and

carboxylated polyvinylidene fluoride.

16. (Original) The method of claim 14, wherein said surface-modified

materials are modified with functional groups selected from the group consisting of

acyl fluoride, esters, amino, carboxyl, hydroxyl, epoxide, thiol, and alkanethiols.

17. (Original) The method of claim 1, wherein said target biopolymer is

immobilized on the substrate by direct adsorption or covalent attachment.

18. (Original) The method of claim 1, wherein said support is in the form of

foams, filaments, threads, sheets, films, slides, gels, membranes, beads, plates, and

planar devices having discrete isolated areas in the form of wells, troughs,

pedestals, hydrophobic or hydrophilic patches, die-cut adhesive reservoirs, or other

physical barriers to fluid flow.

19. (Original) The method of claim 1, wherein the surface of said support is

modified to contain hydrophobic and/or hydrophilic regions prior to said dispensing

step.

20. (Original) The method of claim 1, wherein said substrate is wetted with

an organic modifier selected from the group consisting of ethanol, methanol,

isopropanol, 2-butanol, acetic acid, dextran sulfate and polyacrylic acid, prior to said

dispensing step.

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21. (Previously presented) The method of claim 1, further comprising codispensing an internal standard with said sample to determine the concentration of

said target biopolymer in said aliquots.

22. (Currently amended) The method of claim 1, wherein in step (b), said

microarray is contacted with a plurality of [probes] probe biopolymers.

23. (Currently amended) The method of claim 22, wherein each aliquot is

contacted with a different probe biopolymer.

24. (Currently amended) The method of claim 22, wherein said [probes]

probe biopolymers are labeled with identical reporter groups.

25. (Currently amended) The method of claim 22, wherein said [probes]

probe biopolymers are labeled with reporters that are distinguishable from one

another.

26. (Currently amended) The method of claim 1, wherein in step (b), each of

said aliquots is contacted with a plurality of [probes] probe biopolymers.

27. (Currently amended) The method of claim 26, wherein said [probes]

probe biopolymers are labeled with reporters that are distinguishable from one

another.

28. (Original) The method of claim 1, wherein said aliquots are deposited

onto said substrate at about 1 to 1536 sites per square millimeter of the substrate

surface area.

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29. (Original) The method of claim 1, wherein said substrate is a multiple well microplate, and said aliquots are deposited at between 1 to 1536 sites per well of said microplate.

- 30. (Currently amended) A method for detecting a target nucleic acid in a sample, comprising:
 - (a) preparing a microarray of said sample by dispensing aliquots of said sample at discrete sites onto a substrate and immobilizing said target nucleic acid on said substrate, wherein the microarray is an array of dots, each dot having a diameter from about 1 to 500 microns, wherein each of said aliquots contains the same amount of said target nucleic acid;
 - (b) contacting said microarray with [a] one or a plurality of labeled nucleic acid [probe] probes under hybridizing conditions that allow the formation of [a complex between] one or a plurality of complexes, each complex comprising said target nucleic acid and one of said [probe] probes, wherein said probe is a nucleic acid that is substantially complementary to said target nucleic acid, wherein said labeled nucleic acid [probe is applied to dots individually] probes are deposited on said dots in said microarray; and
 - (c) detecting the presence of <u>and quantifying</u> said [complex] <u>complexes</u> as a measurement for the presence or the amount of said target nucleic acid in said sample.

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31. (Original) The method of claim 30, wherein the preparation of said microarray further comprises dispensing said sample aliquots on said substrate by

a method selected from the group consisting of jet printing, piezoelectric dispensing

methods, solenoid dispensing methods, thermal dispensing methods, solid pin

contact printing methods, capillary quill contact printing methods, microfluidic-

based printing, and silk screening.

32. (Original) The method of claim 30, wherein said aliquots comprise

picomole amounts of said target nucleic acid.

33. (Canceled)

34. (Original) The method of claim 30, wherein said aliquots comprise

femtomole amounts of said target nucleic acid.

35. (Original) The method of claim 30, wherein said aliquots comprise

attomole amounts of said target nucleic acid.

36. (Original) The method of claim 30, wherein said aliquots comprise

zeptomole amounts of said target nucleic acid.

37. (Original) The method of claim 30, wherein said target nucleic acid is

selected from the group consisting of single-stranded RNA, mRNA, single-stranded

DNA, double-stranded DNA, amplified DNA, cDNA and PNA.

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38. (Original) The method of claim 30, wherein said labeled probe is selected

from the group consisting of single-stranded RNA, mRNA, single-stranded DNA,

double-stranded DNA, amplified DNA, cDNA, and PNA.

39. (Original) The method of claim 30, wherein said probe is labeled with a

reporter selected from the group consisting of dyes, chemiluminescent compounds,

enzymes, fluorescent compounds, metal complexes, magnetic particles, biotin,

haptens, radio frequency transmitters, radioluminescent compounds, radioactive-

labeled biomolecules, dye-labeled beads, quantum dots, and bar coded particles.

40. (Original) The method of claim 30, wherein said substrate is made of

crosslinked polymers, porous foam, nitrocellulose, nylon, glass, silica, ceramic, gold,

porous metallic materials, non-porous metallic materials, and surface-modified

materials.

41. (Original) The method of claim 40, wherein said crosslinked polymers are

selected from the group consisting of polypropylene, polyethylene, polystyrene, and

carboxylated polyvinylidene fluoride.

42. (Original) The method of claim 40, wherein said surface-modified

materials are modified with functional groups selected from the group consisting of

acyl fluoride, esters, amino, carboxyl, hydroxyl, epoxide, thiol, and alkanethiols.

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43. (Original) The method of claim 30, wherein said substrate is wetted with an organic modifier selected from the group consisting of ethanol, methanol, isopropanol, 2-butanol, acetic acid, dextran sulfate and polyacrylic acid, prior to said

dispensing step.

44. (Original) The method of claim 30, further comprising co-dispensing an internal standard with said sample to determine the concentration of said target

nucleic acid in said aliquots.

45. (Original) The method of claim 30, wherein in step (b), the microarray is

contacted with a plurality of probes.

46. (Original) The method of claim 45, wherein each aliquot is contacted with

a different probe.

47. (Original) The method of claim 45, wherein each probe is labeled with an

identical reporter.

48. (Original) The method of claim 45, wherein said probes are labeled with

reporters which are distinguishable from one another.

49. (Original) The method of claim 30, wherein each of said aliquots is

contacted with a plurality of probes.

50. (Original) The method of claim 49, wherein said probes are labeled with

reporters which are distinguishable from one another.

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51. (Original) The method of claim 30, wherein said aliquots are deposited

onto said substrate at about 1 to 1536 sites per square millimeter of the substrate

surface area.

52. (Original) The method of claim 30, wherein said substrate is a multiple

well microplate, and said aliquots are deposited at between 1 to 1536 sites per well

of said microplate.

53. (Currently amended) A method for identifying one or more target

analytes in a sample, comprising:

(a) preparing a microarray of said sample by dispensing aliquots of said

sample at discrete sites onto a substrate and immobilizing said

analytes on said substrate, wherein the microarray is an array of dots,

each dot having a diameter from about 1 to 500 microns, wherein each

of said aliquots contains the same amount of said target analytes;

(b) contacting said microarray with a plurality of labeled probes specific

for each of said target analytes under conditions that allow formation

of a [complex between each] plurality of complexes, each complex

comprising one of said target analytes and one of said labeled [probe]

probes specific for said target analyte, wherein said plurality of

labeled probes are [applied to dots individually] deposited on said dots

in said microarray; and

(c) detecting and quantifying said complexes as a measurement of the

presence or the amount of said target analytes.

54. (Original) The method of claim 53, wherein the preparation of said

microarray further comprises dispensing said sample aliquots on said substrate by

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a method selected from the group consisting of jet printing, piezoelectric dispensing

methods, solenoid dispensing methods, thermal dispensing methods, solid pin

contact printing methods, capillary quill contact printing methods, microfluidic-

based printing, and silk screening.

55. (Original) The method of claim 53, wherein said analyte is selected from

the group consisting of biopolymers, drugs, small organic molecules, nucleic acids,

proteins, receptors, antigens, carbohydrates, cells, cellular fragments, and tissues.

56. (Original) The method of claim 53, wherein said probe is selected from

the group consisting of nucleic acids, antibodies, antibody fragments, ligands, and

carbohydrates.

57. (Original) The method of claim 53, wherein said label is selected from the

group consisting of dyes, chemiluminescent compounds, enzymes, fluorescent

compounds, metal complexes, magnetic particles, biotin, haptens, radio frequency

transmitters, radioluminescent compounds, radioactive-labeled biomolecules, dye-

labeled beads, quantum dots, and bar coded particles.

58. (Original) The method of claim 53, wherein said aliquots comprise

picomole amounts of said analyte.

59. (Original) The method of claim 53, wherein said aliquots comprise

femtomole amounts of said analyte.

60. (Original) The method of claim 53, wherein said aliquots comprise

attomole amounts of said analyte.

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61. (Original) The method of claim 53, wherein said aliquots comprise

zeptomole amounts of said analyte.

62. (Original) The method of claim 53, wherein said substrate is made of

crosslinked polymers, porous foam, nitrocellulose, nylon, glass, silica, ceramic, gold,

porous metallic materials, non-porous metallic materials, and surface-modified

materials.

63. (Original) The method of claim 53, wherein said surface-modified

materials are modified with functional groups selected from the group consisting of

acyl fluoride, esters, amino, carboxyl, hydroxyl, epoxide, thiol, and alkanethiols.

64. (Original) The method of claim 53, wherein the surface of said support is

modified to contain hydrophobic and/or hydrophilic regions prior to said dispensing

step.

65. (Original) The method of claim 53, wherein said substrate wetted with an

organic modifier selected from the group consisting of ethanol, methanol,

isopropanol, 2-butanol, acetic acid, dextran sulfate and polyacrylic acid, prior to said

dispensing step.

66. (Original) The method of claim 53, further comprising co-dispensing an

internal standard with said sample to determine the concentration of said analytes

in said aliquots.

67 (Original) The method of claim 53, wherein each aliquot is contacted with

a different probe.

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68. (Original) The method of claim 67, wherein each probe is labeled with an

identical reporter.

69. (Original) The method of claim 67, wherein each probe is labeled with a

different reporter.

70. (Original) The method of claim 53, wherein each aliquot is contacted with

a plurality of probes.

71. (Original) The method of claim 53, wherein said aliquots are deposited

onto said substrate at about 1 to 1536 sites per square millimeter of the substrate

surface area.

72. (Original) The method of claim 53, wherein said substrate is a multiple

well microplate, and said aliquots are deposited at between 1 to 1536 sites per well

of said microplate.